# PATENT SPECIFICATION

Inventor: FRANKLIN VEATCH.

752,345



Date of Application and filing Complete Specification Jan. 7, 1954.

No. 510/54.

Complete Specification Published July 11, 1956.

-Classes 2(2), F2D(1:2), F2G, F3(A:C:E:F), F4; 2(5), P6(A: Index at acceptance: C13A: D1); and 91, F1.

### COMPLETE SPECIFICATION

## A method of Inhibiting the Evaporation of Volatile Products

We, THE STANDARD OIL COMPANY, a Corporation organised under the laws of the State of Ohio, United States of America, of Midland Building, Cleveland, Ohio, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

This invention relates to the inhibition of the evaporation loss of volatile products by a floating layer comprising small hollow hole-

free particles.

Various industries have long been plagued 15 with the problem of evaporation of products stored in tanks. Attempts to minimise this loss have resulted in various proposals, none of which has proved entirely satisfactory. In the petroleum industry, for example, the shape of

20 the storage tanks, particularly the roofs of said tanks have been altered in an effort to reduce the evaporation loss but this has been only a partial solution due to the "breathing" effect with a change in temperature and upon filling

25 and emptying the tank. Floating foams of myriad chemical compositions have been proposed, but with no appreciable reduction in evaporation loss, probably due to the short life of such foams. Furthermore, some of these 30 chemical compositions have contaminated the This evaporation loss has stored products.

proved exceedingly costly.

Accordingly, an object of this invention is the development of a permanent, inexpensive 35 and effective method of substantially decreasing or inhibiting the evaporation loss from the surfaces of products containing volatile com-

This and other objects, as well as the utility 40 and advantages of the invention will become apparent from the following detailed descrip-

It has now been found that the evaporation loss of volatile products can be effectively 45 decreased and inhibited by floating thereon a layer comprising particles formed from film-forming materials which are hollow, predomi-[Price 3s. Od.]

nantly hole-free, have a gas within and are very small. Each particle is a separate, unitary discrete entity.

It has been found that a floating layer of particles in themselves inhibits evaporation of

volatile products.

It has also been found that this inhibition can be markedly improved by the presence of 55 a sealing liquid in order to fill up the capillary interstices between the surfaces of the particles, which sealing liquid may be naturally contained in the volatile product, or may be separately added as an ingredient of the foam. 60 The sealing liquid is held in place by capillary forces between the surfaces of the particles. The sealing liquid in effect is floated or held on top of the volatile product by the particles.

Accordingly the present invention provides 65 a method of inhibiting the evaporation of volatile products which comprises covering the surface of said volatile product with a floating layer of individual small, hollow, gas-filled, hole-free particles of a material inert to said 70 volatile product, which particles have an average diameter of less than 500 microns and a liquid displacement density of from 0.05 to

The present invention also includes a foam 75 floating on a volatile product to inhibit evaporation thereof comprising individual small, hollow, gas-filled, hole-free particles of a material inert to said volatile product, which particles have an average diameter of less than 80 500 microns and a liquid displacement density of from 0.05 to 0.6.

One method of preparing the hollow particles is described in co-pending Application No. 13150/53 (Serial No. 743,866), the disclosure of which (to the extent is describes the particles and method of making them) is

incorporated by reference.

The method of preparing these hollow, gasfilled predominantly hole-free particles, as 90 described in said co-pending application, consists essentially in the steps of spraying a volatile vehicle containing a film-forming material capable of forming a tough gas-impermeable

Price 3s. 64

Price 330

skin, and a gaseous substance or a gas-evolving substance, into a current of heated air whereby the vehicle is volatilised and the filmforming material is solidified in the form of hollow particles containing the evolved gas. The gas evolved during the drying is trapped within the particles and prevents the collapse of the film wall during drying or thereafter.

Gases per se utilisable in the above des-10 cribed process include carbon dioxide, methyl chloride, ammonia, dimethyl ether, ethylene oxide, methylamine, methyl bromide and dimethylamine. Gas-evolving substances, more commonly called blowing agents, are also utilisable in the process. Some of the wellutilisable in the process. known blowing agents which are operable are inorganic and organic salts of carbonates, nitrites, carbamates, oxalates, formates, benzoates, sulphites, and bicarbonates, such as the 20 sodium, ammonium, calcium and magnesium, salts thereof. Organic substances such as phydroxy phenylazides, di-N-nitrosopiperazines, polymethylene nitrosamines such as di-Nnitrosopentamethylene tetramine 25 methylene trinitrosamine, compounds containing two or more groups of the formula CON (alkyl NO, such as succin-bis-(N-nitroso-methylamide)-diazoaminobenzene, diazoiso-

butyric acid dinitrile and homologues thereof 30 prepared using cyclohexanone or methyl ethyl ketone istead of acetone, can also be utilised in the aforedescribed method of preparing hollow, hole-free particles.

Many of the above blowing agents will 35 react with other substances to produce gases instantaneously. Carbonates and sulphites, for example, such as sodium carbonate and sodium sulphite, react with acids such as hydrochloric or sulphuric acid to produce carbon dioxide 40 and sulphur dioxide respectively. Ammonium

salts react with bases such as sodium hydroxide to liberate ammonia. Therefore, by feeding in hydrochloric acid solution to a solution of a film-forming material containing a carbonate

45 or sulphite just as the solution is entering the atomiser, carbon dioxide or sulphur dioxide is liberated and is present during spray-drying as the gas necessary for forming hole-free hollow particles.

50 The amount of gas-forming substance required depends on the weight of the film-forming solution, the type and amount of gas evolved, and is usually within the range of about 0.1% to 25% by weight of the solution 55 to be atomised.

The film-forming materials used may be organic or inorganic. The organic materials include cellulose derivatives, such as cellulose acetate, cellulose acetate-butyrate, and cellulose acetate-propionate, thermoplastic synthetic

60 lose acetate-propionate, thermoplastic synthetic resins, such as polyvinyl resnis, i.e. polyvinyl alcohol, polyvinyl chloride, co-polymers of vinyl chloride and vinyl acetate, polyvinyl butyral, polystyrene, polyvinylidene chloride, 65 acrylic resins, such as polymethyl methacryl-

ate, polyallyl resins, such as polyallyl alcohol and polyallyl acetate, polyethylene, and polyamide (Nylon) resins, and thermosetting resins in the initial water- or organic solvent-soluble stage of partial polymerisation, the resins being 70 converted after or during formation of the particles into a more or less fully polymerised solvent-insoluble stage, such as alkyd, polysiloxane, phenol-formaldehyde, urea-formaldehyde and melamine formaldehyde resins. 25 All these resins are film-forming and therefore capable of forming tough-skinned particles during evaporation of droplets of solutions thereof in aqueous or organic solvents. Natural film-forming materials may 80 also be used including soybean protein, zein protein, alginates, and cellulose in solution as cellulose xanthate or cuprammonium cellulose. Inorganic film-forming substances include sodium silicates, polyborates and polyphos- 85

The solvent employed, will, of course, be dictated by the solubility of the film-forming material used. The solvent should, upon evaporation, be conducive to solidification of 90 the material, so that a tough skin is quickly formed over the surface of the droplet. Water, alcohols, ethers, esters, organic acids, hydrocarbons and chlorinated hydrocarbons, are the

most noteworthy satisfactory solvents.

The concentration of the film-forming material solution is not critical, the lower limit being controlled by the size of the particle since the smallest particles are formed from dilute solutions, and the upper limit being 100 set by the viscosity of the solution and by the maximum particle size desirable. Optimum results have been obtained with solutions containing from 1 to 15% of film-forming material, although concentrations up to 30% have also been found satisfactory. Solutions as dilute as 0.1% have given satisfactory results. Aqueous solutions which contain from 1% to 10% of polyvinyl alcohol (viscosity of a 4% aqueous solution at 20° C. is 4 to 28 cp by the Hoeppler falling-ball method) or from 1% to 30%, preferably 5% to 20%, of phenol-formaldehyde resin, are very satisfactory.

The solution is prepared, sub-divided into droplets, and dried, by conventional means. 115 The use of spray-drying equipment, in which the droplets are dried in a current of hot gas, usually air, is especially advantageous. The drying temperature is adjusted according to the stability and softening point of the film-forming material, the size of the droplets produced and the volatility of the solvent employed. However, as those skilled in the art appreciate, because of the cooling effect of evaporation, drying air of very high temperatures may be used without injury to low melting or easily decomposable materials. A high drying rate is very desirable; usually, air temperatures in the range of 80 to 700° F. will be adequate. Satisfactory drying condi-

752,345

الرودي.

tions for individual cases are shown in the examples.

The dry particles that are produced in conventional spray-drying equipment, employing solutions of the concentrations indicated above, are smaller than can be seen with the naked eye. Frequently their diameter is about ten times the thickness of the plastic skin surrounding their hollow interior, but these 10 dimensions will depend upon the droplet size produced by the equipment used and the concentration of the film-forming material in the solution.

The following Examples I to VI illustrate 15 the preparation of the hollow particles.

EXAMPLE I.

An aqueous solution of a water-soluble partially polymerised phenol-formaldehyde resin (Durez 15281 the word "Durez" being a 20 Registered Trade Mark) was prepared with a blowing agent. The composition contained 2½% of the resin, ½% of di-N-nitrosopentamethylenetetramine on an active basis, and 97% of water. The composition was spray 25 dried at the rate of 57 cc. per minute utilis-ing a laboratory spray drier. The drier operated at an inlet temperature of 680° F. and an outlet temperature of 445° F. The particles obtained had a bulk density of 0.013 gm/cc., 30 and a liquid displacement of 0.070. The average size of the particles was 33.6 microns as determined with a Subsieve Sizer and the range of particle size as determined with a microscope was 5 to 60 microns. In order to 35 determine if particles were hole-free, a weighed quantity of particles was floated in a bath of petroleum naphtha at room temperature and the percentage of the original sample which sank after 24 hours was determined. In this 40 particular example none of the particles sank.

Example II.

Example I was repeated using 10% of the resin, 1% of the blowing agent and 89% of water. The composition was spray dried in a 45 semi-commercial apparatus at the rate of 1 lb. per hour with a drier inlet temperature of 530° F. and an outlet temperature of 425° F. The run was repeated except that the feed rate was increased to 3 lbs. per hour. Inasmuch 50 as the particles obtained at the two different feed rates were not substantially different, the output of the two runs was combined. The paricles had a bulk density of 0.098 gm/cc., a liquid displacement of 0.32, an average (Subsieve Sizer) size of 16 microns, and a microscopic range of 2 to 30 microns. In the flotation test approximately 4% of the particles sank in the naphtha.

Example III.

60 A 10% solution of a phenol-formaldehyde resin (Durez 14798) in water containing 1% of ammonium carbonate as a blowing agent with 89% of water was spray dried at a feed rate of 111 cc. per minute, with a drier operating at an inlet temperature of 700° F. and an 65 outlet temperature of 400° F. In the flotation test only 3% of the particles sank after 24 hours. The particles had a liquid displacement density of 0.449, a bulk density of 0.16, and a size range (microscopic) of 2 to 30 microns. 70

3 .

EXAMPLE IV.

Example III was repeated except that ammonium nitrite was employed as the blowing agent. The particles had a liquid displacement density of 0.353, a bulk density of 0.09, and 75 a microscopic size of 2 to 40 microns. Only 1% sank after twenty-four hours in the flotation test

EXAMPLE V.

A 5% aqueous solution of polyvinyl alcohol (Dupont "Elvanol" Grade 70—05) was prepared with ½% ammonium carbonate as the blowing agent. The balance comprised water in the amount of 94.5%. This composition was spray dried at the feed rate of 80 85 cc. per minute, at an inlet temperature of 600° F. and an outlet temperature of 270° F. The particles had a liquid displacement density of 0.192, a bulk density of 0.07, and a microscopic size range of 5 to 50 microns. Only 2% 90 sank in the flotation test.

EXAMPLE VI.

An aqueous 10% solution of sodium silicate was prepared containing  $2\frac{1}{2}\%$  (1% active) of a mixture of 40% of dinitrosopentamethylenetetramine and 60% of filler and the solution spray dried. The air temperature at the inlet was 700° F. and at the outlet 360° F., and the solution was fed in at a rate of 153.8 cc. per minute. The product had a dry bulk density of 0.236 grams per cc., and a liquid displacement of 0.580. The average particle size was 22.2 microns, as determined by a "Subsieve Sizer", and the range of particle size observed in the microscope from 5 to 40 105 microns.

Following the general description and the examples given heretofore, particles formed from urea-formaldehyde resin, polystyrene, and methyl cellulose were prepared having bulk densities within the range of 0.05 to 0.23, and an average particle size of 2.02 to 36.5 microns.

In addition to hollow particles made by the above described process, it is possible to utilise hollow particles made by other processes in which the wall of the particle can be formed of other materials. For example, the particles may be formed from a ceramic material, such as may be made by heating clay particles at a very high temperature or by the formation of hollow hole-free glass beads. These particles will float on a liquid and can be used alone or in combination with the particles described in greater detail heretofore.

The particles used in the invention include.

all such very small, hollow, predominantly hole-free particles of more or less spherical shape which will float on a liquid, irrespective of the material from which they are formed 5 and the manner in which they are made as long as they are insoluble in and inert with respect to the materials with which they are in contact, and irrespective of the process by which they are made.

The preferred bulk density of the particles is within the range of 0.01 to 0.30. The liquid displacement density is within the range of 0.05 to 0.6 (gm/cc) preferably 0.2 to 0.5. The bulk density is of interest when transporting 15 or storing the particles. The liquid displacement density is of interest when the space between the particles is to be occupied by a liquid such as in a foam. The particles have an average diameter of 1 to 500 microns, gener-

20 ally 25 to 250 microns. Particles having a size range of less than 1 micron are entirely suitable but are more difficult to collect following

the spray-drying process.

The sealing liquids, whether contained in 25 the volatile product or added to the particles, must possess inherent properties of non-volatility, chemical stability, non-corrosiveness, and non-solvency for the particles. When an added sealing liquid is employed it should pre-

30 ferably be insoluble in the volatile product. These properties are all physical characteristics which are known. It can readily be ascertained what liquids have these properties. The co-action between the particles and the sealing 35 liquid does not involve any chemical reaction.

One skilled in the art, in view of this disclosure, will be able to select liquids having suitable physical properties.

The amount of sealing liquid, when one is 40 employed, that is desirable in the formation of the particle suspension is usually within the range of about 40 to 80% by volume of the suspension, 55 to 70% being preferred, depending of course on the nature and amount

45 of particles and on the specific liquid used. If the amount is too small it will not fill the spaces between the particles. If it is too large, there will be more liquid than can fill the

spaces between the particles and the excess 50 will not be floated by the particles. The density of the mixture of the particles and the sealing liquid must be appreciably less than the density of the volatile product to be protected. Consiedring the average or combined

55 density of the sealing liquid and the particles, it is possible to select the liquid and the proportions so that the mixture has a density less than that of the product to be covered, so that the mixture will float. If the product has a

60 higher density, particles of a higher density can be used. The selection and the amount ofthe sealing liquid, and its physical co-action with the particles will be readily apparent to one skilled in the art in view of the disclosure 65 herein.

Any suitable method of preparing this floating foam of particles may be employed. When a sealing liquid is not used or where it is naturally present in the volatile product, it is sufficient to place the particles on top of the 70 product or mix them with it whereupon they will float to the top. When a separate sealing liquid is used, the simplest method consists in forming a suspension of the particles in the sealing liquid by suitable agitation. This suspension or slurry may then be pumped onto the surface of the product to be protected, where it forms a floating protective layer.

The particle content of the foam, when a

sealing liquid is used, must be sufficient to 80 impart buoyancy to the foam and usually constitutes from about 20 to 60%, and preferably 30 to 45% by volume of the suspension. The viscosity of the slurry must be sufficiently low to render it fluid enough for pumping or for 85 other means of installation on the surface of

the volatile product.

A foam layer of about ½ inch to 1 inch in thickness has been found to give adequate protection against evaporation and contamination, 90 although a thickness as little as 1 inch gives some improvement. Greater thickness than 1 inch, while not inoperative, does not give sufficiently superior results to be justified economically. In general, it is not contem-95 plated that a thickness in excess of 3 inches would be used.

The volatile products which may be protected in accordance with the invention are numerous. Included are volatile hydrocarbons 100 such as gasoline, naphthas and other light fractions, including liquefied hydrocarbons such as liquid butane. Alcohols such as methyl, ethyl, and isopropyl alcohols are volatile and may have their evaporation reduced in accord- 105 ance with the invention. Chlorinated hydrocarbons such as trichloroethylene may similarly be protected. Other low boiling liquefied gases such as chlorine, liquid oxygen or liquid nitrogen may be stored utilising the inven- 110 tion. Aqueous solutions such as sugar solutions or electroplating solutions may be protected in accordance with the invention. Acetone and low boiling ketones and ethers may have their evaporation reduced by utilising the 115

The sealing liquid, if one is employed, may be selected in relation to the volatile product to be protected. If the volatile product naturally contains a sealing liquid, such as a non-120 volatile oil in a volatile hydrocarbon, the nonvolatile oil will function as the sealing liquid. Where a separate sealing liquid is used such as in protecting a volatile hydrocarbon distillate, the glycols have proved particularly satis- 125 factory, the specific glycol used being dependent on the nature of the substratum to be protected, such as hexylene glycol being particularly useful on crude oil tanks and dipropylene glycol being preferred on tanks 130

storing gasoline and other light materials. Examples of other suitable sealing liquids include ethylene glycol, propylene glycol, polypropylene glycol, higher glycol polymers, polymers of olefin oxides such as ethylene and propylene oxides (available as Ucon oils), castor oil, Carbitol, Methylcellosolve, hexylene glycol (2-methyl-2,4-pentane diol), dimethyl phthalate, triethylene tetramine, tetraethylene pentamine, and polyvinyl methyl ether.

The alcohols, aqueous solutions and similar materials may be protected with a hydrocarbon oil such as white oil, which is substantially inert. Acetone and ethers and the liquefied gases may be protected with a perfluoro oil of the general formula C<sub>n</sub>F<sub>2n+2</sub>, where n is an

integer of 12 or more.

In selecting a separate sealing liquid it is desirable to employ one having a relatively high 20 viscosity. This enhances the retention of the sealing liquid in the capillary interstices between the particles. If the sealing liquid has too low a viscosity there is a tendency for the buoyant action of the particles to squeeze the 25 sealing liquid out of the capillary spaces. This tends to form a crust at the top, rendering resealing more difficult if the floating foam layer On the other hand, the viscosity is broken. should not be too high or the foam will become so heavy and viscous as to be relatively immobile, rendering it difficult to spread over the surface and secure an effective seal. The choice of sealing liquids and viscosities thereof will be obvious from the disclosure 35 herein and the specific examples, depending upon the size and buoyancy of the particles, the substratum to be sealed, the length of life desired in the foam, and other factors apparent to one skilled in the art from the disclosure herein. When a separate sealing liquid is employed, hollow particles of small size, such 40 herein.

as under 75 microns, are preferred.

The following specific examples are illustrative of foams that may be employed in prac-

45 tising the invention.

### EXAMPLE VII.

A floating layer of particles made in accordance with Example I, two inches thick, was floated on liquid butane, and the evaporation 50 was substantially reduced. This contains no natural or added sealing liquid.

### EXAMPLE VIII.

A floating layer of particles made in accordance with Example 1 was applied in a layer 55 one inch thick on a wide cut catalytic cracker feed stock boiling between 150° and 800° F. The heavier components in this stock accumulate in the space between the particles and form a sealing liquid which is naturally present in the volatile product.

Example IX.

A floating layer especially adapted for use on gasoline is as follows:—

| % by vol. Particles (formed from phenol- formeldehyde regin liquid dis-   | 65  |
|---|-----|
| formaldehyde resin, liquid displacement density: 0.070) - 41.5  Sealing liquid (composed of 70% of dipropylene glycol and 30% of high viscosity polymeric ole-  | 70  |
| fin oxide (Ucon oil 75—H— 90000) 58.5   | 70  |
| A floating layer especially adapted for cut-back asphalt is as follows:—  % by vol.   | 75  |
| Particles (formed from polyvinyl alcohol, liquid displacement density: 0.19) 28.2 Scaling liquid (castor oil) 71.8  | 80  |
| EXAMPLE XI.  A floating layer for use on gasoline has the   |     |
| following composition:  % by vol.  Particles (formed from urea-form- aldehyde, liquid displacement  | 85  |
| aldehyde, liquid displacement density: 0.35) 46.7 Sealing liquid (polyvinyl methyl ether) 53.3  |     |
| Example XII. % by vol.  | 90  |
| Particles (formed from phenol- formaldehyde, liquid displace- ment density: 0.2) 40  Sealing liquid (heavy medicinal white oil, the material sold under the Registered Trade Mark "Nujol") 60                                   | 95  |
| Mark "Nujol") 60 This foam is especially adapted for use on alcohols and aqueous solutions.  EXAMPLE XIII.  | 100 |
| Particles (formed from urea-<br>formaldehyde resin, liquid dis-<br>placement density: 0.15) - 35<br>Perfluoro oil (C <sub>16</sub> F <sub>38</sub> , viscosity<br>about 60 millipoises and essen-                               | 105 |
| tially non-volatile at 100° C.) - 65 This foam is useful on chlorinated hydrocarbons, chemicals, such as ketones, and ethers in which the particles are insoluble. Other agents may be added to the foam to                     | 110 |
| evaporation loss from storage tanks containing<br>non-volatile products, such as rust inhibitors  | 115 |
| fire-resistant agents, anti-bleeding agents, etc. Our Specification No. 13151/53 (Serial No. 744,070) claims a method of inhibiting the evaporation of volatile components of crude oil which comprises covering the surface of | 120 |

oil which comprises covering the surface of said crude oil with a floating layer of small,

hollow, hole-free particles and crude oil or a

non-volatile component of crude oil, said hollow particles having an average particle diameter of from 1 to 500 microns, a liquid displacement density of 0.05 to 0.6 gm./cc., and 5 occupying from 20 to 70% by volume of the floating layer, and we make no claim herein to anything claimed in Specification No. 13151/53.

Subject to the foregoing disclaimer, what

10 we claim is:

1. A method of inhibiting the evaporation of volatile products which comprises covering the surface of said volatile product with a floating layer of individual small, hollow, gas-15 filled, hole-free particles of a material inert to said volatile product, which particles have an average diameter of less than 500 microns and a liquid displacement density of from 0.05 to

2. A method as claimed in claim 1 in which the volatile product is a volatile liquid petro-

leum product.

3. A method of inhibiting the evaporation of volatile products which comprises covering 25 the surface of said volatile product with a floating layer of a foam containing as the essential ingredients a non-volatile sealing liquid and individual small, hollow, gas-filled hole-free particles of a material inert to said volatile 30 product and said sealing liquid, which particles have an average diameter of less than 500 microns and a liquid displacement density of from 0.05 and 0.6.

4. A method as claimed in claim 3 in which 35 the sealing liquid is naturally present in the

volatile product.

5. A method as claimed in claim 3 in which the sealing liquid is not naturally present in the volatile product and is insoluble therein.

6. A method as claimed in any one of the preceding claims 3 to 5 in which the volatile

product is a volatile liquid petroleum hydro-

carbon.

7. A foam floating on a volatile product to inhibit evaporation thereof comprising indivi- 45 dual small, hollow, gas-filled, hole-free particles of a material inert to said volatile product, which particles have an average diameter of less than 500 microns and a liquid displacement density of from 0.05 to 0.6.

8. A foam floating on a volatile product to inhibit evaporation thereof comprising a non-volatile sealing liquid and individual small, hollow, gas-filled, hole-free particles of a material inert to said volatile product, which 55 particles have an average diameter of less than 500 microns and a liquid displacement density of from 0.05 to 0.6, the sealing liquid in said foam being present in an amount within the range of 40 to 80% by volume and the par-60 ticles in said foam being present in an amount within the range of 20 to 60% by volume to impart a density to said foam less than the density of volatile product and to hold the sealing liquid in the interstices between the 65 particles.

9. A foam as claimed in claim 8 in which the sealing liquid is naturally contained in the

volatile product.

10. A foam as claimed in claim 8 in which 70 the sealing liquid is not contained in the volatile product and is insoluble therein.

11. A method of inhibiting the evaporation of volatile products substantially as described with reference to any one of Examples VII to 75

**ELKINGTON & FIFE** Consulting Chemists and Chartered Patents Agents, Bank Chambers, 329, High Holborn, London, W.C.1, Agents for the Applicants.

Leamington Spa: Printed for Her Majesty's Stationery Office, by the Courier Press.—1956. Published at the Patent Office, 25, Southampton Buildings, London, W.C.2, from which copies may be obtained.